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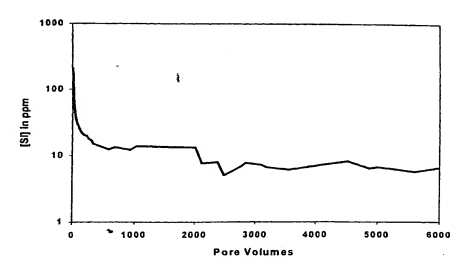
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(54) Title: MICROCAPSULE WELL TREATMENT

Scale Inhibitor Release Profile from a Plastic Capsule



(57) Abstract: A material and a device for releasing chemicals in a fluid environment is disclosed, and a method for releasing substances into downhole fluid environments. The invention also relates to materials for sand control and hydraulic fracturing. The substance release device typically comprises a polymeric material for releasing a substance into a downhole fluid environment in a well, and in preferred embodiments the material comprises a spherical, plastic chemical release capsule which is hard, permeable and may encapsulate a range of solids and/or liquids for subsequent release. These solids or liquids can include inhibitors such as scale inhibitors and other oilfield production chemicals for release into the wellbore of an oil or gas producing well or a water injection well.

01/94744 A1

WO 01/94744 A1



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microcapsule well treatment

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2	
3	This invention relates to material for releasing
4	chemicals in a fluid environment, and to a method of
5	producing the material. In particular, this invention
6	relates to materials for sand control and hydraulic
7	fracturing, the same materials releasing chemicals in
8	a fluid environment, and to a method of producing
.9	these materials.
10	•
11	Advances in drilling and completion technology have
12	revolutionised new field development and in-fill
13	drilling strategies. The use of subsea and/or
14	horizontal well completions is now common and in some
15	cases these complex wells require sand control
16	measures to be installed to maintain the integrity of
17	the wellbore and prevent it collapsing. A variety of
18	well completions are conventionally used for sand
19	control purposes including fracture packing and
20	gravel-packing techniques, prep-packed screens, wire

WO 01/94744 PCT/GB01/02482

. 1	wrapped screens and expandable screens. It is
2	preferred that the flow of fluids through the sand
3	control system is not impaired since this can reduce
4	the productivity and/or injectivity of the well.
5	Hydraulic fracturing is used to improve well
6	productivity. It is therefore preferred that the flow
7	of fluids into and through the propped fracture is
8	not impaired since this can also reduce the
9	productivity and/or injectivity of the well.
10	
11	The installation of subsea and/or horizontal well
12	completions also increase the complexity, logistical
13	difficulty and cost of intervention for the
14	mitigation of production chemistry problems such as
15	inorganic scale formation, asphlatene and wax
16	deposition, corrosion and bacterial growth on the
17	wellbore conduits.
18	
19	According to the present invention there is provided
20	a substance release device comprising a polymeric
21	material for releasing a substance into a downhole
22	fluid environment in a well.
23	i
24	In preferred embodiments the material comprises a
25	spherical, plastic chemical release capsule which is
26	hard, permeable and may encapsulate a range of solids
27	and/or liquids for subsequent release. These solids
28	or liquids can include inhibitors such as scale
29	inhibitors and other oilfield production chemicals
30	for release into the wellbore of an oil or gas
31	producing well or a water injection well.

1	The material can be used for sand control and for
2	hydraulic fracturing (for clarity the material shall
3	herein be termed "a proppant" when it is applied to
4	hydraulic fractures) and well clean up. The material
5	can be used for sand control purposes whilst also
6	providing controlled release of oilfield production
7	chemicals from within the sand control medium. This
8	includes application in gravel packed, fracture
9	packed and pre-packed screen sand control systems.
10	The material can also be used for more effective well
11	clean up after completion in both injection and
12	production wells via the controlled release of enzyme
13	based materials that release acids to dissolve any
14	mineral deposits and breakdown residual bio-polymers
15	from the well completion process. The material can
16	be used for hydraulic fracturing purposes whilst also
17	providing controlled release of oilfield production
18	chemicals from within the proppant medium. In
19	addition this material may be used for hydraulic
20	fracturing and either or both the controlled release
21	of oilfield chemicals and proppant flow back control.
22	The material may also be used within the rathole of a
23	well to provide controlled release of an oilfield
24	chemical into the aqueous environment.
25	
26	Preferably the polymeric material is used for sand
27	control and hydraulic fracturing.

29 Preferably the polymeric material is permeable.

The material can comprise a polymeric plastics 1 material such as polypropylene, polyethylene, high 2 density polyethylene, high density polypropylene, 3 polyethylene terephthalates, polyamides (both 4 aliphatic and aromatic), liquid crystal polymers, 5 liquid engineered resins, starch and polyhydroxy 6 alkanoates or mixtures of these. 7 8 The material typically comprises a matrix that 9 provides the material with support, and which 10 contains the chemical to be released. 11 12 13 Other materials may be used as the matrix such as acrylic, polybutylene, polycarbonate, polyester, 14 15 polystyrene, polyurethane, polyvinyl chloride, polycaprolactone, polybutylene terephthalate, 16 polyvinyl alcohol, polylactic acid, polyglycolide, 17 polyester amide, polyimides, acrylonitrile-butadiene-18 19 styrene, acrylonitrile-styrene-acrylate, 20 polyoxymethylene, polybutylene, polyisobutylene, 21 polyvinylbutyral, epichlorohydrin elastomer, nitrile 22 elastomer, nitrile rubber, polyetherketone, polyetheretherketone; polyetherketoneketone, 23 polymethylmethacrylate, polyethylene oxide, 24 polyphenylene oxide, polysulphones, 25 polyethersulphone, polyurea, chlorinated 26 27 polyethylene, ethylene-chlorofluoroethylene, 28 tetrafluoroethylene-perfluoropropylene, 29 Perfluoroalkoxy, silicon rubbers and other polymeric 30 materials consisting of mixtures of the above and 31 copolymers, terpolymers and

1	hydrophobically/hydrophilically modified and
2	crosslinked derivatives of the above.
3	
4	Other materials can be used instead of or in addition
-5	to the polymeric or plastics material, such as
6	thermoplastic rubber, resins, hot melt adhesives,
7	fibreglass, silicones, fluorosilicones e.g. Ryton,
8	polysiloxanes, fluoroelastomers e.g. Viton, Aflas,
9	fluorocarbons e.g. PTFE, PVDF, halogenated polymers,
10	cellulose, polysaccharides, lignin, chitin, gums and
11	mixtures and derivatives of the above.
12	
13	In addition waxes, calcium stearate, and metallocene
14	can be used instead of or in addition to the
15	polymeric or plastics material.
16	
17	The material contains a high active content of scale
18	inhibitor, or the chemical to be released, typically,
19	10-90wt%/v. The high mass of inhibitor contained
20	within the capsule can reduce (and on some wells
21	potentially eliminate) the need for scale related
22	intervention. In addition, the high loading of
23	inhibitor offers the potential for significant
24	treatment lifetime with minimal product deployment.
25	
26	The release of the scale inhibitor can be further
27	controlled by changing its solubility in the brine
28	phase or by restricting contact between the inhibitor
29	salt and the brine phase. The solubility of the
30	inhibitor in the brine phase can be reduced by
31	increasing the ratio of the polyvalent cation to

1	inhibitor in the precipitated salt or by changing the
2	nature of the polyvalent cation. Careful selection
3	of the polymer type or the use of hydrophobic
4	additives can reduce water ingress into the porous
5	matrix and thus reduce inhibitor release.
6	
7	In certain embodiments the hardness of the material
8	can be varied by changing the chemical composition of
9	the polymer plastic matrix and/or cross-linking the
10	polymer plastic material or by reinforcing the
11	particle matrix with natural, synthetic, glass or
12	metal fibres, sand, silicates or other minerals and
13	ceramic matrices such as bauxite, resins, carbon or
14	boron based materials and impact modifiers. Typically
15	the unconfined compressive strength would be in the
16	range 500-100,000psi.
17	
18	In other embodiments the material can be selected to
19	have an oil tolerance which can be adjusted by
20	changing the chemical composition of the polymer
21	plastic matrix and/or modifying the hydrophobic or
22	hydrophilic properties of the polymer matrix or
23	cross-linking the polymer matrix with a suitable
24	reagent. In addition, the chemical resistance to
25	acids and oil may be increased by the addition of a
26	chemical stabiliser, for example Ebonite.
27	
28	The capsules are typically of permeable polymeric
29	material, and are manufactured using an underwater
30	pelletising system consisting of an extrusion and
31	either a spheronisation or granulation process. The

. 1	capsules typically contain a high active content of
2	scale inhibitor, typically, 10-90wt%/v. The
3	particles can be produced over a wide range of sizes.
4	The size distribution of the particles in a sample of
5	material can be large, with a heterogeneous
6	distribution of particles of different sizes, or
7	small, with a fairly homogeneous range of similarly
8	sized particles (so called mono dispersed particles).
9	The range of preferred sizes is typically $250\mu m$ -
10	5mm, with the distribution of size being typically
11	±5% of the mean size.
12	
13	The density of the capsules can be controlled by the
14	variation of the type of polymer matrix and/or by
15	including a weighting agent such as barite, zirconium
16	oxide, manganese oxide, titanium dioxide, tungsten or
17	magnetite. The hardness, compressive strength and
18	elastic properties (rigidity and
19	elongation/deformation) can be controlled by the
20	variation of the type of polymer matrix and/or
21	including a strengthening agent such as glass, sand,
22	minerals, carbon or boron fibres and/or an impact
23	modifier. The unconfined compressive strength is
24	typically between 500-100,000psi.
25	•
26	The material is typically extruded from a pelletising
27	system. A single or twin screw system can be used,
28	and other mixers such as Farrell and Banbury mixers
29	or a Feeder extruder can be used.
30	

Mono dispersed spherical particles (i.e. a relatively 1 2 homogeneous mixture of similarly sized particles) can 3 be produced over a wide range of particle size. 4 typically 250µm - 5mm, with the distribution of size 5 being typically ±5% of the mean size. The variation б in particle size can allow certain embodiments of the 7 invention to provide gravel packs of different mesh 8 sizes, where in certain embodiments there is an 9 advantage in having a narrow range of particle sizes 10 in the one application. The spherical and mono dispersed nature of the particles typically provides 11 better packing in the gravel pack and a more uniform 12 13 pore throat size. This can provide more effective 14 sand control and reduce the turbulence in the pore 15 throats, thus improving the gravel pack permeability. 16 17 In certain embodiments, the material can have a low 18 and adjustable density (specific gravity usually between 0.9 and 2.0 where mostly polymeric materials 19 20 are used, and even higher SG values up to 12 can be 21 achieved with a higher proportion of heavier materials e.g. mostly tungsten) and that can be 22 23 adjusted by changing the chemical composition of 24 polymeric matrix and/or by adding a weighting agent 25 such as barite, tungsten, zirconium oxide or 26 magnetite or any other material denser than the 27 polymer. The optional low density of the material 28 can provide more effective gravel pack placement and 29 simplify the gravel-packing operation, especially in 30 deviated wells because the high viscosity fluids 31 normally needed to carry dense proppant and/or gravel

pack material through a pipeline system will not be 1 2 needed. 3 In certain embodiments, typically for application in 4 5 rat holes, the polymer matrix can slowly degrade over 6 a time period of 1 week to 10 years leaving little or This will typically enable re-treatment 7 no residue. 8 of the product into the rathole once the product has degraded. The lifetime of the plastic capsule can be 9 adjusted by changing the chemical composition and 10 11 molecular weight of the polymeric matrix and/or 12 incorporating a dissolution agent such as heavy aromatic napthas, peroxides, per sulphates, enzymes 13 14 and metal oxide catalysts into the spherical plastic 15 capsule. The rate and lifetime of release of the encapsulated chemical can be controlled such that it 16 17 corresponds to the degradation lifetime of the 18 polymer matrix material. This may be achieved by 19 varying the amount of encapsulated chemical in the 20 polymer matrix material, changing the chemical 21 composition of the polymer matrix and the inhibitor 22 salt or by coating the polymer matrix and/or coating 23 the encapsulated inhibitor salt. 24 25 In certain embodiments, for application in gravel 26 packs and hydraulic fractures, the polymeric material 27 is preferably somewhat pliable so that it can display 28 a slight amount of deformation at the point of 29 contact, increasing the surface area of contact

between particles. In gravel packs and hydraulic fractures this reduces the effect of applied stress

1 and can result in improved sand and proppant flowback 2 control respectively. The amount of deformation of 3 the polymer matrix required to control sand 4 production and proppant flowback can be adjusted by 5 changing the chemical composition of the matrix 6 and/or including a strengthening agent such as glass, sand, minerals, carbon or boron fibres and/or an 7 impact medifier. Laboratory data from stressed 8 9 cyclic flow back tests with mixtures of conventional 10 proppant and chemically impregnated conventional 11 porous proppant had indicated that although these mixtures passed the initial conductivity tests the 12 13 proppant pack collapsed under cyclic stress and 14 flowed out of the test apparatus. However, when the 15 same tests were repeated with mixtures of 16 conventional proppant and this polymeric material no 17 proppant flow back was observed. This was 18 attributed to cohesion of the proppant pack induced 19 by the flowback control characteristics of the 20 polymeric material. 21 22 Most preferred embodiments of the material comprise 23 spherical capsules although other shapes may be in 24 accordance with the invention. Spherical capsules 25 have the additional advantage for gravel pack 26 applications in that they can be packed more 27 efficiently into a defined space than other shapes of 28 material. In addition, the use of a more homogeneous 29 particle size typically leads to the formation of 30 more homogeneous pore throats. This can afford the 31 gravel pack more effective sand control

1 characteristics and can reduce the turbulence within 2 the sand or proppant pack, which may lead to an increased effective permeability. 3 4 5 In preferred embodiments of the invention the 6 material is charged with any scale inhibitor suitable 7 for the purpose of inhibiting scale in a well bore This may include water soluble organic 8 formation. 9 molecules with carboxylic acid, aspartic acid, maleic 10 acids, sulphonic acids, phosphonic acid and phosphate .11 esters groups including copolymers, ter-polymers, 12 grafted copolymers and derivatives of the above. 13 Examples of such compounds include aliphatic 14 phosphonic acids such as diethylene triamine penta 15 (methylene phosphonate) and polymeric species such as polyvinylsulphonate. The scale inhibitor can be in 16 17 the form of the free acid but is preferably in the 18 form of mono and polyvalent cation salts such as Na, K, Al, Fe, Ca, Mg, NH4. 19 20 21 The chemicals can be solid and mixed with the matrix 22 to give a generally homogeneous mixture comprising 23 the plastics matrix and the chemical. Alternatively, 24 the scale inhibitors etc can be encapsulated in 25 liquid form e.g. by being immobilised, absorbed or 26 encapsulated in an appropriate matrix such as a 27 diatomised clay, resin, starch or other 28 polysaccharide material, polyvinyl alcohol and super-29 adsorbing polymers and then being incorporated in 30 solid form into the main product matrix.

The material can contain scale inhibitor in most 1 2 preferred embodiments but instead in other 3 embodiments either no chemical is included in the formulation or a wide range of other oil field 4 5 chemicals such as inhibitors can be carried including 6 but not limited to corrosion inhibitors, biocides, 7 wax and asphaltene control chemicals, pour point suppressants, dispersants, hydrate and halite 8 inhibitors, demulsifiers, gel breakers, tracers, drag 9 10 reducers and well clean up chemicals including These can also be encapsulated in either 11 enzymes. solid or liquid form into the material. 12 13 Certain embodiments of the present invention tackle 14 sand and combined sand/scale control in gravel packed 15 16 and pre-packed screen wells by using a hard, 17 permeable, spherical plastic capsule, optionally 18 carrying a chemical substance such as an inhibitor, 19 although other embodiments can simply provide an 20 alternative material to sand or gravel for use in 21 gravelpacks. The plastic capsule can be manufactured 22 using a hot melt extrusion and spheronising process 23 and can optionally contain approximately 10-90wt% of active scale inhibitor and can benefit from a low and 24 25 adjustable density. The high mass of inhibitor 26 contained within the particle can reduce the need for 27 scale-related intervention and offers the potential 28 for significant treatment lifetime. The low density 29 of the particle also aids placement and simplifies 30 the gravel packing operation. In addition, the 31 spherical nature and mono dispersed characteristics

1	of the plastic capsule can provide more effective
2	packing and a uniform pore throat size for the gravel
3	pack. This can provide improved sand control and
4	increase the pack permeability through reduced
.5	turbulence at pore throats. The provision of a
6	material for sand and/or combined sand/scale control
7	in e.g. gravel packed and pre-packed screen wells
8	comprising a hard, optionally permeable material with
9	or without a loading of chemical to be delivered to
10	the well is another aspect of the present invention,
11	typically where the material comprises a number of
12	particles having a narrow size range e.g. 400um+/-5-
13	10%.
14	
15	The invention also provides a method of treating a
16	well, the method comprising loading a substance into
17	a polymeric material, inserting the material into the
18	well, and allowing the substance to leach from the
19	polymeric material into the well.
20	
21	Typically the material is inserted into a fissure,
22	fracture, screen area, gravel pack, fracture pack or
23	a pre-packed screen, usually as part of the well
24	completion.
25	
26	The invention also provides a method of forming a
27	chemical-releasing particle for release of chemicals
28	in a fluid environment, the method comprising mixing
29	the chemical to be released with a matrix material
30	forming the particle, and thereafter forming the
31	particle (e.g. by melting, mixing and/or extrusion of

WO 01/94744 PCT/GB01/02482

14

1 the mixture) from the mixture of the matrix and the 2 chemical, so that the chemical is dispersed 3 (preferably homogeneously) throughout the formed 4 particle. 5 6 In an extruder system, the polymeric matrix material 7 and the other additives are typically fed into the top of the extruder via a hopper at the rear of the 8 9 extruder system. The extruder is preferably an auger 10 type mixer, which is typically heated to melt the 11 polymer matrix. The auger screw system then 12 typically mixes and disperses the other additives in 13 the molten polymer as the mixture typically moves 14 along the auger screw. Typically a vent is provided 15 before the material reaches the head of the extruder 16 to allow gas or moisture to escape. However this 17 venting can be further enhanced by fitting a vacuum 18 pump to more effectively withdraw gas and moisture. 19 The mixture typically proceeds into a throat which 20 typically slows down the mixture and typically 21 increases the pressure before typically releasing the mixture in the form of strands of typically circular 22 23 cross section that are extruded through apertures in 24 an extrusion head. The strands are then typically 25 cooled in a water bath and typically pelletised into 26 cylindrical pellets using a chopper. The temperature 27 of the heated extruder screw is normally dependent on 28 the melting point of the polymers, but the typical 29 extrusion temperatures vary between 100°C and 400°C.

. 1	The invention also provides a polymeric downhole
2	proppant.
3	
4	The invention also provides a method of supporting a
5	well or a portion thereof, comprising disposing a
6	support material in the well or portion to be
7	supported, wherein the support material comprises a
8	polymeric material.
9	
10	The invention also provides a polymeric material for
11	use downhole for the control of particulate matter
12	entering the production stream of the well.
13	
14	The invention also provides a method of controlling
15	the entry of particulate matter into the production
16	stream of a well, the method comprising disposing a
17	number of polymeric devices into the wellbore between
18	the formation and the production fluids outlet of the
19	well.
20	
21	Examples of the present invention will now be
22	described by way of illustration only and with
23	reference to the accompanying figures, wherein
24	
25	Fig. 1 is a schematic representation of
26	apparatus for producing devices according to the
27	invention;
28	Fig. 2 is a graph plotting the concentration of
29	scale inhibitor against the number of cycles of
30	fluid passing through a closed system;

WO 01/94744 PCT/GB01/02482

1	Figure 3 shows the plot of compressive stress
2	(psi) versus axial strain for certain
3	embodiments of devices;
4	Figure 4 shows a graph of pore throat size
5	distribution versus gravel pack compressive
6	stress as determined from a mercury stress
7	injection test on a further embodiment;
8	Fig 5 shows a thin section photomicrograph of a
9	sand slurry injection test of a further
10	embodiment;
11	Fig 6 shows a graph of the % weight loss of
12	total material and scale inhibitor versus the
13	degradation time of the another embodiment of
14	material in days at 107°C; and
15	Fig 7 shows scale inhibitor release profile
16	under simulated rathole conditions for PHA
17	polymer in produced brine at 107°C in another
18	embodiment.
19	·
20	To form the capsules an underwater pelletising
21	process is used, which utilises an extruder having
22	either a single screw or twin screw system. A single
23	screw extruder comprises a steel tube of variable
24	length, which has one single shaft screw that rotates
25	at speed in the tube as it is heated, melting the
26	polymer and mixing it as it is pushed down the tube.
27	A twin screw extruder consists of two co-rotating
28	screws in a single barrel with a variety of mixing
29	zones down the length of the heated barrel in which
30	both screws work together to mix and blend. This
31	works the material more effectively compared with a

. 1	single screw extruder and a better dispersion of the
2	solid additives into the polymer is achieved.
3	
4	The dispersion of the material additives into the
5	binder or polymer can be divided into four steps:
6	
7	(a) wetting of the additives = pre-mixing of the
8	additives with the binder;
9	(b) dispersion of agglomerates into aggregates and
10	primary particles;
11	(c) distribution into the binder or polymer;
12	(d) stabilisation against re-agglomeration.
13	
14	The plastic/chemical mix is then introduced into a
15	known underwater pelletising system shown in Fig. 1.
16	The Fig. 1 system is suitable generally for producing
17	pellets according to the invention.
18	
19	The pre-extruded cylindrical pellets or the polymeric
20	matrix material and additives for example inhibitors
21	and weighting agents are fed via a hopper into an
22	extruder or melt pumps which force the molten mixture
23	through a screen changer and/or polymer diverter
24	valve, which helps build up the pressure at the head,
25	and for the start-up of the run. A gear pump can be
26	used to regulate the flow of polymer to the head to
27	keep the pressure evenly distributed throughout the
28	system. The mixture then flows through a die 10
29	having a series of holes arranged in a circular
30	pattern. As the mixture emerges from the holes, it
31	is cut into lengths by rotating blades 12 and is

1	solidified into pellets by process water which is
2	pumped through conduits 15 across the material face
3	in the cutting chamber 17. Tempered water transfers
4	the pellets via conduits 20 to a centrifugal dryer
5	and spheroniser 22 where the water and the pellets
6	are separated and the dry spherical pellets are
7	discharged via chute 25. The process water is
8	filtered, pressurised, tempered and returned to the
9	Cutting chamber 17. The system can produce pellets
10	(e.g. mono dispersed pellets) typically at between 10
11	to 6000kg/hr in a size range typically from 40µm to
12	5mm ±5%.
13	
14	The extrusion process may be combined with a Farrell
15	bridge consisting of two mixing areas. In the first
16	area the mixture is melted and mixed in an upper
17	chamber using single screw, before it is dropped down
18	to two co-rotating screws in the bottom chamber,
19	where further mixing of the product occurs before it
20	is delivered to the head of the chamber. A venting
21	process takes place between the top and bottom
22	chambers to remove any moisture. This venting
23	process can be enhanced by use of a vacuum pump
24	system.
25	
26	A hole may be cut into the side of the extruder,
27	usually two-thirds of the way down the barrel and it
28	is used to feed additives and fillers directly into
29	the barrel, commonly termed a feeder extruder.
30	

31

An oil-heated drum mixer or Banbury Mixer may be used 1 to slowly turn and blend the polymers and additives. 2 The additives usually need to be more thermally 3 stable and the mixing process is much slower. 5 The spherical plastic capsule may have a variable oil 6 7 tolerance which can be adjusted by changing the 8 chemical composition of the polymer plastic matrix 9 and/or modifying the hydrophobic or hydrophilic 10 properties of the polymer plastic matrix or cross-11 linking the polymer plastic matrix with a suitable 12 reagent. To modify the polymer a thermally stable hydrophobic group such as an aliphatic or an aromatic 13 14 amide, lauryl or phosphate ester, or any other 15 hydrophobic group is grafted onto the polymer backbone as a side chain. This provides the polymer 16 17 with improved oil or water resistance depending on 18 the nature of the polymer matrix and the modification 19 In the cross-linking process, the polymer 20 molecules are linked using peroxide, silanes and 21 metallocene chemicals to form a more complex 22 macromolecular structure. This is then cured at 23 various temperatures to achieve improved strength and 24 oil, water, acid and bacterial resistance. 25 26 The scale inhibitors may be encapsulated in liquid 27 form if they are immobilised on an appropriate matrix 28 such as a diatomised clay, resin, super-adsorbing 29 polymer and high porosity versions (~40-50%) of the

process a liquid is incorporated into a solid matrix

In the immobilisation

main polymer matrix material.

1	by a spray drying, shear mixing, emulsion
2	polymerisation, coating or by a simple soaking
3	process in which the liquid is either imbibed into or
4	adsorbed onto the surface of the solid material.
5	
6	Coatings can be applied to the outer surfaces of the
7	capsules; for example, organic surfactants and waxes
8	can be added to the polymer melt or applied to the
9	outer surfaces of the resultant particle to achieve
10	improved oil, water, acid and bacterial resistance.
11	In addition, the polymer melt additive or surface
12	coating can also be used to control the release rates
13	of the encapsulated chemical into the wellbore of an
14	oil or gas producer well or a water injection well.
15	
16	The resultant spherical particles may be injected
17	into an oil well or water injection well. Certain
18	embodiments of the particles can withstand
19	particularly high pressures associated with such
20	wells. The particles may be injected into the rat-
21	hole, hydraulic fractures or into the annulus in
22	gravel packed, fracture packed and pre-packed sand
23	screen completed wells. The chemicals such as scale
24	inhibitors will gradually over time leak out from the
25	particles and so treat the local environment
26	accordingly. The release rate can be controlled by
27	varying the amount of encapsulated chemical in the
28	polymer matrix material, changing the chemical
29	composition of the polymer matrix and the inhibitor
30	salt or by coating the polymer matrix and/or coating
31	the encapsulated inhibitor salt.

Examples · 1 2 Example 1 - Extrusion Temperature 165°C - 200°C 3 4. 5 Particles were produced using the above Fig. 1 device 6 and having the following proportion of constituents:-7 40% PP/HDPE (60/40 mix of polypropylene/high density 8 9 polyethylene) - as the plastic matrix material; The 10 material can be added as a fine powder of average 11 size $0-100\mu m$. 12 13 30% Magnetite (Iron oxide) - as the weighting agent; 14 The material can be added as a fine powder with an 15 average particle size of 1-100µm. 16 17 30% Solid Polycarboxylate - as a scale inhibitor 18 The material can optionally be added as a fine 19 powder with an average particle size of 1-100µm, and 20 typically around 70 µm or less. 21 22 The plastic matrix material and the other two 23 components were added! into the hopper and fed into 24 the top of the extruder. The mixture was then melted 25 within the heated auger type mixer and the components 26 dispersed in the polymer matrix, as described earlier 27 in the manufacturing process, before being extruded 28 using the underwater pelletising system also 29 described earlier. The finished capsule was 30 particularly suitable for rat hole applications where

a weighting agent is normally necessary to keep it in 1 2 place and prevent it from floating out of position. 3 The specific gravity of this material was ~1.4 and 4 it's unsinfined compressive strength was >500psi. 5 The product was also thermally stable in both brine and oil up 140°C. However, above this temperature 6 7 the product will soften and agglomerate and may break 8 down in crude oil. The scale inhibitor was an optional constituent, and this can be omitted in 9 10 embodiments intended for gravelpack applications. 11 12 Example 2 - Extrusion Temperature 165°C - 200°C 13 14 Particles were produced using the above Fig. 1 device 15 and having the following proportion of constituents:-16 17 60% PP/HDFE (60/40 mix of polypropylene/high density 18 polyethylene) - as the plastic matrix material; The 19 material can be added as a fine powder with an 20 average particle size of 1-100µm. 21 22 2% Tungsten - as the weighting agent; The material 23 can be added as a fine powder with an average 24 particle size of 1-100µm. 25 26 38% Solid Polycarboxylate. The material can be added 27 as a fine powder with an average particle size of 1-28 100µm. 29 30 This capsule was manufactured as detailed for the 31 previous example. The product is typically suitable

1	for scale control in rat holes. The reduction in the
2	amount of weighting agent in the plastic capsule
3	optionally allows more scale inhibitor to be loaded
4	into the product agent, thus potentially increasing
5	the treatment lifetime. The reduction in the amount
6	of weighting agent also increases the compressive
7	strength. The specific gravity of this material was
8	~1.20 and its unconfined compressive strength was
9	greater than 1000psi. The product displayed a
10	thermal stability similar to that in Example 1.
11	Again, the scale inhibitor was an optional
12	constituent, and this can be omitted in embodiments
13	intended for gravelpack applications.
14	
15	Example 3 - Extrusion Temperature 200-220°C
16	
17	Particles were produced as described for the previous
18	2 examples and having the following proportion of
19	constituents:-
20	
.21	50% PP/HDPE (60/40 mix of polypropylene/high density
22	polyethylene) - as the plastic matrix material. The
23	material can be added as a fine powder with an
24	average particle size of 1-100μm
25	
26	30% Glass - Strengthening Agent. The material can be
27	added as a fine powder with an average particle size
28	of 1-100μm.
29	•
30	20% Solid THPS (Tetrakishydroxymethylphosphonium

_	barphate, as a brocker sait. The material can be
2	added as an agglomerate with an average particle size
3	of 50-2000μm.
4	
5	This capsule was manufactured as detailed for the
6	previous example. The extrusion temperature was
7	slighter higher to ensure the glass which was added
8	as a fine powder was properly dispersed during the
9	extrusion process. The addition of glass strengthens
10	the capsule by up to 10% as well as increasing the
11	oil resistance and thermal stability of the product
12	to over 150°C. The specific gravity of this material
13	was ~1.20. The unconfined compressive strength was
14	1100psi. This capsule can be used for the control of
15	sulphate reducing bacteria to prevent the generation
16	of H_2S and associated corrosion problems.
17	
18	Example 4 - Extrusion Temperature 200°C - 240°C
19	
20	Particles were produced as described above for the
21	previous three examples and having the following
22	proportion of constituents:
23	· · · · · · · · · · · · · · · · · · ·
24	80% PET/PP (20/80 mix of polyethylene
25	terephthalate[PET)/polypropylene[PP]) - as the
26	plastic matrix material; the material can be added
27	as a fine powder with an average particle size of 1-
28	100µm.
29	

20% Solid Phosphonate - as a scale inhibitor salt. 1 The material can be added as a fine powder with an 2 average particle size of 1-100µm. 3 4 5 This capsule was manufactured as detailed for the previous two examples. The inclusion of polyethylene 6 7 terephthalate in the main matrix material 8 strengthened the bead by a factor of 3 and can be 9 suitable for applications that require the capsule to 10 withstand high pressures. In addition, the inclusion 11 of PET imparts more oil resistance and increase the 12 thermal stability of the product to >180°C. 13 specific gravity of the material was ~1.3 and it's 14 unconfined compressive strength was >3400psi. 15 16 Example 5 - Extrusion Temperature 200 - 260°C 17 18 This capsule was manufactured as detailed for the 19 previous two examples and having 100% Polyamide 20 (Nylon) as the plastic material. 21 22 The use of polyamide as the main matrix material 23 provided even more strength than the bead as quoted 24 in Example 4. The compressive strength was again 25 increased by a factor of 3 and use of Polyamide increases the chemical resistance and thermal 26 27 stability of the product to >250°C. The material can 28 be suitable for sand control in applications that 29 require the plastic capsule to withstand extremely 30 high pressures and, in addition, the polyamide matrix 31 material can display a slight amount of deformation

.1 and can be used for proppant flowback control in 2 hydraulic fractures and gravel packs. Typically this embodiment excludes any substance to be released from 3 4 the capsule, but can incorporate such substances if 5 desired. 6 7 The specific gravity of this material was ~1.2-1.3 8 and its unconfined compressive strength exceeded 9 12000psi. 10 Example 6 - Extrusion Temperature 200°C -260°C 11 12 13 This capsule was manufactured as detailed for the 14 previous two examples and has the following 15 proportion of constituents: 16 17 70% Polyamide - as the plastic matrix material. The 18 material can be added as a fine powder with an average particle size of 1-100μm. 19 20 21 30% Solid Phosphonate - as a scale inhibitor salt. 22 The material can be added as a fine powder with an 23 average particle size of 1-100 um. 24 25 This capsule can be used for sand and proppant flow 26 back control and combined sand/proppant flow back and 27 scale control in gravel packs, fracture packs, screen 28 wells and hydraulic fractures. 29 30 Fig. 2 shows the concentration of scale inhibitor 31 released versus the number of pore volumes of fluid

1	oluted through the packed material. One many relieve
. 1	eluted through the packed material. One pore volume
2	is defined here as approximately 25% of the total
3	volume of the packed material and corresponds to the
4	volume in the system that is not occupied by the
5	plastic material. It can be seen from Fig. 2 that
6.	the release of scale inhibitor is constant even after
7	6000 pore volumes. Therefore, the scale inhibitor is
8	released at a constant low level for long periods of
9	time.
10	
11	It can be seen from Fig. 3 that the material in
12	Example 5 does become slightly strained as the stress
13	is increased to > 8000-10000psi. Therefore a slight
14	amount of deformation of the material does occur in
15	both the presence and absence of scale inhibitor.
16	
17	The specific gravity of this material was ~ 1.3 and
18	its unconfined compressive strength was greater than
19	10,500psi.
20	•
21	Example 7 - Extrusion Temperature 165°C
22	
23	This capsule was manufactured as detailed for the
24	previous two examples and having the following
25	proportion of constituents:-
26	
27	40% HDPE (high-density polyethylene) - as the plastic
28	matrix material. The material can be added as a fine
29	powder with an average particle size of 1-100μm.
30	· · · · · · · · · · · · · · · · · · ·

Т.	30% Barite (Darium sulphate) - as the weighting
2	agent. The material can be added as a fine powder
3	with an average particle size of 1-500μm.
4	
5	30% solid polycarboxylate - as a scale inhibitor
6	salt. The material can be added as a fine powder with
7	an average particle size of 1-100 µm.
8	
9	The specific gravity of this material was ~1.3 and
10	its compressive strength was >600psi. The product was
11	also thermally stable in both brine and oil up to
12	between 140°C. The finished capsule is particularly
13	suitable for scale control in rat hole applications.
14	
15	Example 8 - Extrusion Temperature 165°C
16	
17	This capsule was manufactured as detailed for the
18	previous two examples and having the following
19	proportion of constituents:-
20	
21	40% HDPE (high-density polyethylene) - as the plastic
22	matrix material. The material can be added as a fine
23	powder with an average particle size of 1-100 µm.
24	
25	30% Rutile (titanium dioxide) - as the weighting
26	agent. The material can be added as a fine powder
27	with an average particle size of 1-500μm.
28	

. 1	30% solid polycarboxylate - as a scale inhibitor
2	salt. The material can be added as a fine powder
3	with an average particle size of 1-100μm.
4	
5	The specific gravity of this material was ~ 1.3 and
6	its compressive strength was >1000psi. The product
7	displayed similar thermally stability to material in
8	example 7. The finished capsule is particularly
9	suitable for scale control in rat hole applications.
10	
11	Example 9 - Extrusion Temperature 200°C -260°C
12	
13	This capsule was manufactured as detailed for the
14	previous two examples and having the following
15	proportion of constituents:-
16	
17	65% Polyamide/HDPE (74/16 mix of polyamide/high
18	density polyethylene) - as the plastic matrix
19	material; The material can be added as a fine powder
20	with an average particle size of $1-100\mu m$.
21	
22	28% Solid Phosphonate - as a scale inhibitor salt.
23	The material can be added as a fine powder with an
24	average particle size of 1-100µm.
25	
26	2% lauric diethanol amine - as an anti static agent.
27	This material can either be added as a liquid or as a
28	fine powder with an average particle size of 1-100µm.
29	

1 5% Calcium Stearate - as a lubricant. The material can be added as a fine powder with an average 2 3 particle size of 1-100µm. 4 5 This capsule can be used for sand and proppant flow 6 back control and combined sand/proppant flow back and 7 scale control in gravel packs, fracture packs, screen wells and hydraulic fractures. 8 9 10 Figure 4 shows the pore throat size distribution 11 versus the gravel pack compressive stress as 12 determined from a mercury stress injection test (as 13 documented in SPE 8294. It can be seen from Fig. 4 14 that the pore throat size distribution is reduced 15 exponentially with increasing compressive stress, 16 with significant deformation occurring at >3000psi, 17 namely, an 87% reduction in pore throat size and a substantial reduction in permeability to air. 18 19 data indicates that at a typical gravel pack 20 compressive stress of 2000psi the product 21 demonstrated acceptable strength characteristics. 22 23 This material can be manufactured down to 500 ± 25um. 24 The specific gravity of this material is ~1.2 and 25 it's confined compressive strength was >2000psi. 26 27 Example 10 - Extrusion Temperature 165°C - 200°C. 28 29 This capsule was manufactured as detailed for the 30 previous two examples and having the following 31 proportion of constituents:-

1	65% HDPE - as the plastic matrix material; The
2	material can be added as a fine powder with an
3	average particle size of 1-100µm.
4	
5	30% Solid Phosphonate - as a scale inhibitor salt.
6	The material can be added as a fine powder with an
. 7	average particle size of 1-50µm.
8	•
9	2% lauric diethanol amine - as an anti static agent.
10	This material can be added as a liquid or as a fine
11	powder with an average particle size of 1-100μm.
12	
13	3% wax - as a lubricant. This material can be added
14	as a molten liquid or as a fine powder with an
15	average particle size of 1-100 μ m.
16	
17	Figure 5 shows a photomicrograph of a thin section of
18	a gravel pack of the above 300µm material taken from
19	a sand slurry injection test. The sand material was
20	injected as a 1wt%/v slurry at 50ml/minute and the
21	average diameter (D50) of the sand particles was
22	100μm. It can be seen from Fig. 5 that none of
2.3	injected sand passed tinto or out of the 300µm gravel
24	pack material. Therefore the above material provided

effective sand control at 300μm.

WO 01/94744 PCT/GB01/02482

1	This material can be manufactured down to 300 \pm 15 μ m.
2	The specific gravity of this material is ~1.2 and it
3	is thermally stable in both oil and brine to >140°C.
4	
5	Example 11 - Extrusion Temperature 180°C -220°C.
. 6	
7	This capsule was manufactured as detailed for the
8	previous two examples and having the following
9	proportion of constituents:-
10	
11	70% PolyHydroxyAlkanoate (PHA) - as the degradable
12	polymer matrix material. This material can be added
13	as an agglomerate with an average granule size of 1-3
14	mm.
15	
16	30% solid phosphonate - as a scale inhibitor salt.
17	This material can be added as a fine powder with an
18	average particle size of 1-100 μm .
19	
20	The finished capsule is particularly suitable for
21	scale control in rat hole applications. The material
22	has been designed to degrade or slowly dissolve so
23	that it is possible to re-treat the rat hole without
24	the need for expensive remedial cleaning procedures.
25	
26	Fig. 6 shows the % weight loss of the total material
27	and scale inhibitor versus the degradation time in
28	days at 107C. It can be seen from Fig. 6 that the
29	material has undergone a substantial total weight
30	loss. A comparison of the total weight loss with the
31	% mass loss profile due to the release of the

encapsulated scale inhibitor, which is relatively 1 constant at a low level, indicated that 2 degradation/dissolution of the polymer matrix was 3 4 occurring. The trends in the data indicate a predicted degradation lifetime of between 12-14 5 months at 107°C. 6 7 Figure 7 shows the concentration of scale inhibitor 8 released versus the number of cell volumes of brine 9 10 eluted at 107°C. The test was performed in a specially designed cell to simulate the release of 11 scale inhibitor from a product placed in the rat 12 hole. It can be seen from Fig. 7 that the PHA 13 polymer matrix displays favourable scale inhibitor 14 release characteristics, with the scale inhibitor 15 16 being released at low levels over a long period. 17 18 The specific gravity of this material was ~1.2-1.3 19 and its compressive strength was >1000psi. The 20 product was also thermally stable in both brine and oil up to between 170-180°C. 21 22 23 Example 12 - Extrusion Temperature 200-220°C 24 25 This capsule was manufactured as detailed for the 26 previous two examples and having the following 27 proportion of constituents:-28 29 61% polyethylene terephthalate(PET) - as the polymer 30 matrix material. This material can be added as a fine 31 powder with an average particle size of 1-100µm.

30% solid phosphonate - as a scale inhibitor salt.

2 This material can be added as a fine powder with an 3 average particle size of 1-100µm. 4 5 9% titanium oxide oxidation catalyst - as a 6 degradation additive. This material can be added as 7 a fine powder with an average particle size of 1-8 100µm. 9 10 The finished capsule is particularly suitable for scale control in rat hole applications. 11 The material 12 has been designed to degrade or slowly dissolve over 13 10-14 months. 14 15 Example 13 - Extrusion Temperature 165-200°C 16 17 This capsule was manufactured as detailed for the 18 previous two examples and having the following 19 proportion of constituents:-20 21 55% starch - as the polymer matrix material. This 22 material can be added as a fine powder with an 23 average particle size of 1-100 mm. 24 25 35% solid potassium hexacyanoferrate III - as a halite inhibitor salt. This material can be added as 26 27 a fine powder with an average particle size of 50-28 500µm. 29

10% Tungsten polyamide blend- as a weighting agent. 1 2 This material can be added as a granule in masterbatch format with an average granule size of 3 4 500-3000μm. 5 6 The finished capsule is particularly suitable for halite scale control in rat hole applications. The 7 8 specific gravity of this material was ~1.1-1.3 and the material is thermally stable up to 140°C. 9 10 11 Example 14 - Extrusion Temperature 165-200°C 12 13 This capsule was manufactured as detailed for the 14 previous two examples and having the following proportion of constituents:-15 16 17 70% polyamide - as the polymer matrix material. This 18 material can be added as a fine powder with an 19 average particle size of 1-100µm. 20 21 30% solid enzyme - as the acid generating enzyme. 22 This material can be added as a granule with an 23 average particle size of 100-1000 µm. 24 25 This material is suitable for sand control and improved well clean up after completion in both 26 . 27 injection and production wells. 28 29 Modifications and improvements may be incorporated. 30 without departing from the scope of the invention.

31

1	Cla	ims
2		
3	1.	A substance release device comprising a
4		polymeric material for releasing a substance
5		into a downhole fluid environment in a well.
6		
7	2.	A device according to claim 1, wherein the
8		substance comprises one or more selected from
9		the group comprising scale inhibitors, hydrate
10		and halite inhibitors, corrosion inhibitors,
11		biocides, wax and asphaltene control substances,
12	•	demulsifiers, gel breakers, tracers, drag
13		reducers and well clean up substances including
14		enzymes; organic molecules, acids, esters, and
15		aliphatic substances.
16		
17	3.	A device as claimed in any preceding claim
18		wherein the polymeric material is permeable.
19		
20	4.	A device as claimed in any preceding claim,
21		wherein the material is one or more selected
22		from the group comprising polypropylene,
23		polyethylene, high density polyethylene, high
24		density polypropylene, polyethylene
25		terephthalates, polyamides (both aliphatic and
26	•.*	aromatic), liquid crystal polymers, liquid
27		engineered resins, starch and polyhydroxy
28		alkanoates or mixtures of these.
29		
30	5.	A device as claimed in any preceding claim,

wherein the material comprises a matrix that

1		provides the material with structural support,
2		and which contains the substance to be released.
3		
4	6.	A device as claimed in claim 5, wherein the
5		material also contains one or more of the group
6		of substances comprising acrylic, polybutylene,
7		polycarbonate, polyester, polystyrene,
8		polyurethane, polyvinyl chloride,
9		polycaprolactone, polybutylene terephthalate,
10		polyvinyl alcohol, polylactic acid,
11		polyglycolide, polyester amide, polyimides,
12		acrylonitrile-butadiene-styrene, acrylonitrile-
13		styrene-acrylate, polyoxymethylene,
14		polybutylene, polyisobutylene, polyvinylbutyral,
15		epichlorohydrin elastomer, nitrile elastomer,
16		nitrile rubber, polyetherketone,
17		polyetheretherketone, polyetherketoneketone,
18		polymethylmethacrylate, polyethylene oxide,
19		polyphenylene oxide, polysulphones,
20		polyethersulphone, polyurea, chlorinated
21		polyethylene, ethylene-chlorofluoroethylene,
22		tetrafluoroethylene-perfluoropropylene,
23		Perfluoroalkoxy, silicon rubbers and other
24		polymeric materials consisting of mixtures of
25		these, copolymers, terpolymers and
26	:	hydrophobically/hydrophilically modified and
27		crosslinked derivatives of these.
28		
29	7.	A device as claimed in any preceding claim,
30		wherein the material contains one or more of the
31		group of substances comprising waxes, calcium

1		stearate, metallocene, thermoplastic rubber,
2		resins, hot melt adhesives, fibreglass,
3		silicones, fluorosilicones e.g. Ryton,
4		polysiloxanes, fluoroelastomers e.g. Viton,
5		Aflas, fluorocarbons e.g. PTFE, PVDF,
6		halogenated polymers, cellulose,
7		.polysaccharides, lignin, chitin, gums and
8		mixtures and derivatives of the above.
9		
10	8.	A device as claimed in any preceding claim,
11		comprising a generally spherical capsule.
12		
13	9.	A device as claimed in any preceding claim,
14		manufactured using an underwater pelletising
15		system using an extrusion and either a
16		spheronisation or granulation process.
17		
18	10.	A device as claimed in any preceding claim,
19		wherein the material contains a high active
20		content of substance to be released.
21		
22	11.	A device as claimed in any preceding claim,
23		including a weighting agent to adjust the
24		density of device.
25		
26	12.	A device as claimed in claim 11, wherein the
27		weighting agent is selected from the group
28		comprising barite, zirconium oxide, manganese
29		oxide, titanium dioxide, tungsten and magnetite
30		

	10.	A device as claimed in any preceding claim,
2		including a strengthening agent to adjust the
3		hardness, compressive strength and/or elastic
· 4		properties of the device.
5		
6	14.	A device as claimed in claim 13, wherein the
7		strengthening agent is selected from the group
8.		comprising glass, sand, minerals, carbon fibres,
9		boron fibres and impact modifiers.
10		
11	15.	A device as claimed in any preceding claim,
12		wherein the material has a specific gravity
13		between 0.9 and 2.0.
14		
15	16.	A device as claimed in any one of claims 1-14,
16		wherein the device has a specific gravity up to
17		12.
18		
19	17.	A device as claimed in any preceding claim,
20		wherein the device and/or the material is
21		pliable.
22		
23	18.	A method of treating a well, the method
24		comprising loading a substance into a polymeric
25		material, inserting the material into the well,
26	,**	and allowing the substance to leach from the
2 7 .		polymeric material into the well.
28		
29	19.	A method as claimed in claim 18, wherein the
30		substance comprises one or more selected from
31		the group comprising scale inhibitors, corrosion

1		inhibitors, hydrate and halite inhibitors,
2		biocides, wax and asphaltene control substances,
3		demulsifiers, gel breakers, tracers, drag
4		reducers and well clean up substances, enzymes,
5		organic molecules, acids, esters, copolymers,
6		ter-polymers, aliphatic compounds, polymeric
7		species, and other oilfield production
8		chemicals.
9		
10	20.	A method according to claim 18 or claim 19,
11		wherein the material releases well cleanup
12		substances.
13		
14	21.	A method as claimed in any one of claims 18-20,
15		wherein the material is inserted into a fissure,
16		fracture, screen area, gravel pack, fracture
17		pack or a pre-packed screen.
18		
19	22.	A method as claimed in any one of claims 18-21,
20		wherein the material is injected into the
21		rathole or base of a well.
22	_	
23	23.	A method as claimed in any one of claims 18-22,
24		wherein the material is used to support portions
25		of the well, reservoir or formation in addition
26	.*	to releasing the substance.
27		
28	24.	A method according to any one of claims 18-23,
29		wherein the material controls entry of
30		particulate matter into the production stream of
31		the well.

25.	A method as claimed in any one of claims 18-24,
	wherein the material is loaded with a high
	active content of the substance to be released.
26.	A method as claimed in any one of claims 18-25,
	wherein the solubility of the substance is
	selected to suit prevailing well conditions.
27.	A method as claimed in any one of claims 18-26,
	wherein the hardness of the material is adjusted
	to suit prevailing well conditions.
28.	A method as claimed in any one of claims 18-27,
	wherein the oil and/or acid tolerance of the
	material is adjusted to suit prevailing well
	conditions.
29.	A method as claimed in any one of claims 18-28,
	wherein a number of the devices having a
	generally heterogeneous size distribution are
	delivered to the well.
30.	A method as claimed in any one of claims 18-28,
	wherein a number of the devices having a
	generally homogeneous size distribution are
, di	delivered to the well.
31.	A method as claimed in claim 30, wherein the
	size distribution of devices delivered to the
	well is ±5% of the mean size.
	26. 27. 28.

1	32.	A method as claimed in any one of claims 29-31,
.2		wherein the range of sizes of the devices
3		delivered to the well is 250µm - 5mm.
4		
5	33.	A method as claimed in any one of claims 18-32,
6		wherein the material is adapted to degrade over
7		a period of time, after which the well is re-
8		treated.
9		
10	34.	A method as claimed in claim 33, wherein the
11		period of degradation is adjusted to suit the
12		prevailing well conditions.
13		
14	35.	A method as claimed in any one of claims 18-34,
15		wherein the rate and lifetime of release of the
16		substance is adjusted to suit the degradation
17		lifetime of the material.
18		
19	36.	A method as claimed in any one of claims 18-35,
20		wherein the material is charged with one or more
21		substances selected from the group comprising
22		scale inhibitors, hydrate and halite inhibitors,
23		corrosion inhibitors, biocides, wax and
24		asphaltene control substances, demulsifiers, gel
25		breakers, tracers, drag reducers and well clean
26	, · ·	up substances including enzymes; organic
27		molecules, acids, esters, and aliphatic
28		substances.
29		·
30	37.	A method of forming a substance-releasing
31		particle for release of substances into a fluid

1		environment in an well, the method comprising		
2		mixing the substance to be released with a		
3		matrix material forming the particle, and		
4		thereafter forming the particle from the mixture		
5		of the matrix and the substance, so that the		
6		substance is dispersed throughout the formed		
7		particle.		
8				
9	38.	A polymeric downhole proppant.		
10				
11	39.	A method of supporting a well or a portion		
12		thereof, comprising disposing a support material		
13	•	in the well or portion to be supported, wherein		
14		the support material comprises a polymeric		
15		material.		
16				
17	40.	A polymeric material for use downhole for the		
18		control of particulate matter entering the		
19		production stream of the well.		
20				
21	41.	A method of controlling the entry of particulate		
22		matter into the production stream of a well, the		
23		method comprising disposing a number of		
24		polymeric devices into the wellbore between the		
25		formation and the production fluids outlet of		
26	. <i>:</i>	the well.		

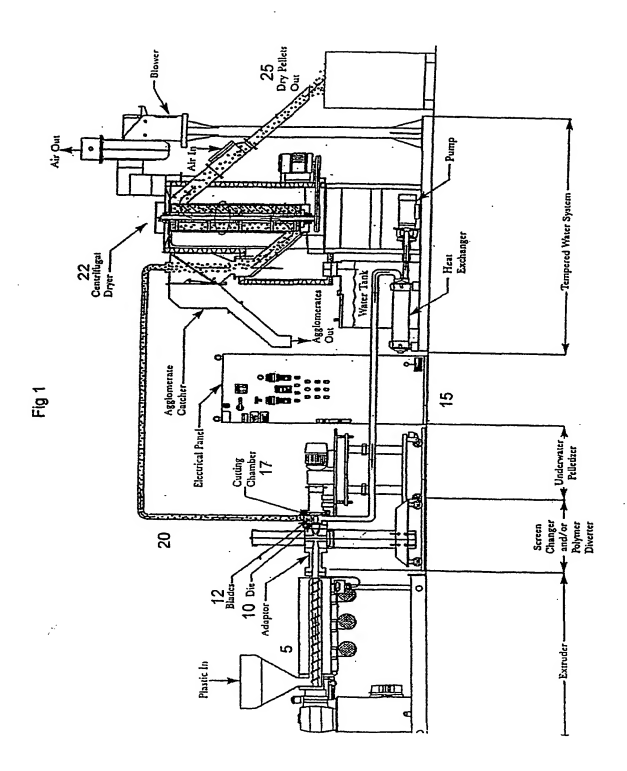
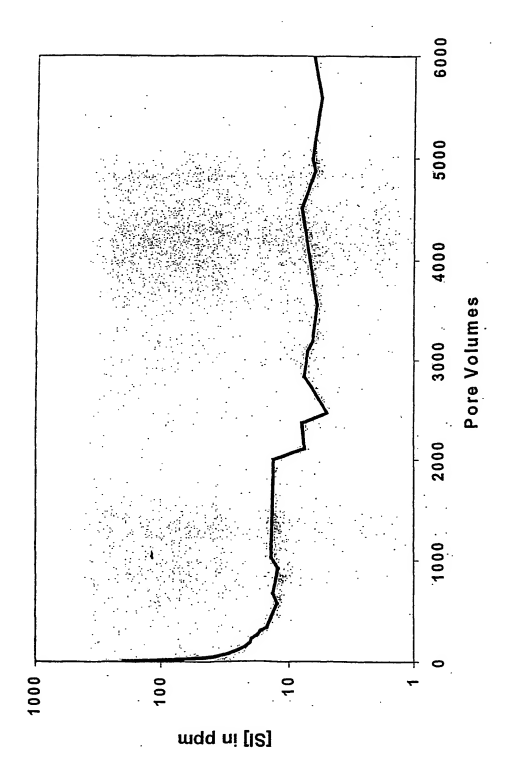


Fig 2 - Scale Inhibitor Release Profile from a Plastic Capsule



···· ··· ···

Fig 3

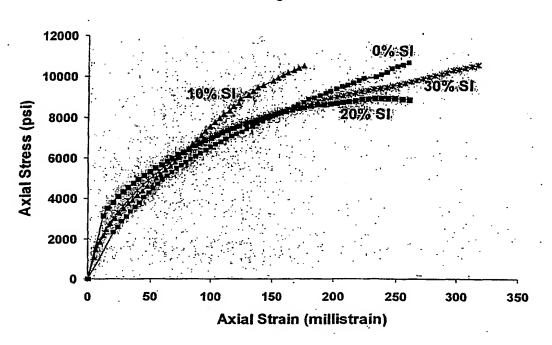
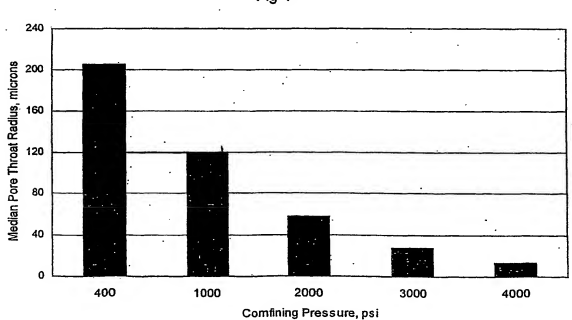


Fig 4



4/5

Fig 5

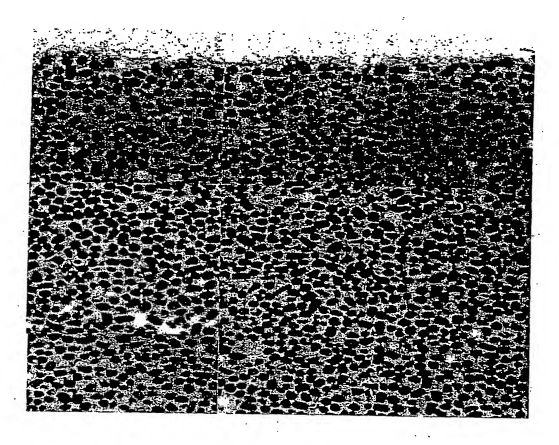
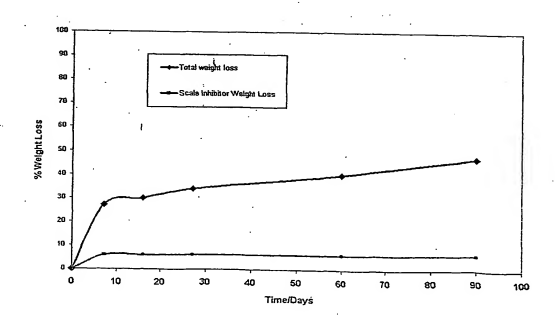
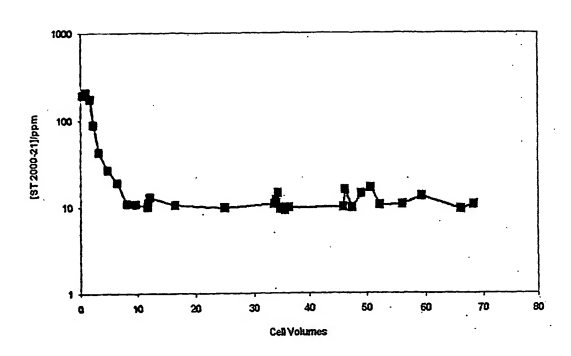


Fig 6



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Fig 7



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INTERNATIONAL SEARCH REPORT

Int Ional Application No PCT/01/02482

A. CLASSIFICATION OF SUBJECT MATTE IPC 7 E21B37/06 E21B43/25

E21B41/02

E21B43/267

According to International Patent Classification	(IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) $IPC\ 7\ E21B\ B01J$

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)

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Special categories of cited documents: A' document defining the general state of the art which is not considered to be of particular relevance E' earlier document but published on or after the international filing date L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) O' document referring to an oral disclosure, use, exhibition or other means P' document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to hvolve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the actual completion of the International search 27 September 2001	Date of mailing of the international search report $04/10/2001$
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Rampelmann, K

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